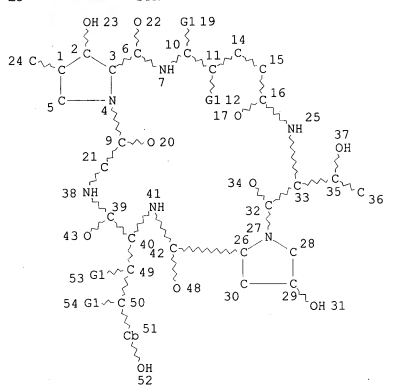
09/673836

(PILE PREGISTRY ENTERED AT 12:14:05 ON 17 OCT 2002) STR

L5



VAR G1=H/OH NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS UNS AT 51 DEFAULT ECLEVEL IS LIMITED

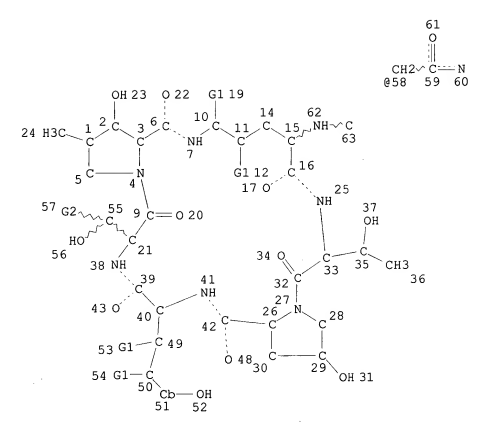
GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 47

STEREO ATTRIBUTES: NONE

L7 1919 SEA FILE=REGISTRY SSS FUL L5

L22

STR



VAR G1=H/OH VAR G2=H/CH3/58 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS UNS AT 51 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RSPEC I

NUMBER OF NODES IS 56

STEREO ATTRIBUTES: NONE

L23 1861 SEA FILE=REGISTRY SUB=L7 SSS FUL L22

L24 314 SEA FILE=REGISTRY ABB=ON PLU=ON L23 AND NR=4

(FINE HEAPLUS! ENTERED AT 12:23:39 ON 17 OCT 2002) L25 99 S L24/P

FILE 'REGISTRY' ENTERED AT 12:29:56 ON 17 OCT 2002 E "C4-HOMOTYROSINE"/CN 5 E "C4-HTYR"/CN 5

FILE 'HCAPLUS' ENTERED AT 12:30:23 ON 17 OCT 2002

1 S L25 AND (C4(W) (HTYR OR (H OR HOMO) (W) (TYR OR TYROSINE))

(FILE 'REGISTRY' ENTERED AT 12:37:53 ON 17 OCT 2002) * See last pgs. for E RANEY NICKEL/CN 5 term C4-homotyrosme

L39

1 S E3

FILE 'HCAPLUS' ENTERED AT 12:37:58 ON 17 OCT 2002

2 S L25 AND (L39 OR RANEY(W)(NICKEL OR NI))

=> s 126 or 140

1641 2 16216 OR 140°

=> sel hit 141 1-2 rn E1 THROUGH E7 ASSIGNED

=> d 1-2 ibib abs hitstr

L41 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:117192 HCAPLUS

DOCUMENT NUMBER:

132:165211

TITLE: INVENTOR(S):

SOURCE:

Method for the production of an antibiotic agent Connors, Neal C.; Petersen, Leslie A.; Hughes, David L.; Dimichele, Lisa M.; Novak, Thomas J.

PATENT ASSIGNEE(S):

Merck & Co., Inc., USA PCT Int. Appl., 18 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA!	rent i	NO.		KII	ND I	DATE			A	PPLI	CATI	ON NO	ο.	DATE		
WO	2000															
	W:	ΑE,	AL,	AM,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CN,	CR,	CU,	CZ,
		EE,	GD,	GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KG,	KR,	ΚZ,	LC,	LK,
		LR,	LT,	LV,	MD,	MG,	MK,	MN,	MX,	NO,	ΝZ,	PL,	RO,	RU,	SG,	SI,
		SK,	TJ,	TM,	TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA,	AM,	ΑZ,	BY,	KG,
		KZ,	MD,	RU,	ТJ,	TM										
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
		DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
			CG,													
AU	9953													19990	0804	
EP	1100	947		A.	1 :	2001	0523		E	P 19	99-9:	3893	3	19990	0804	
														NL,		MC,
			ΙE,													
PRIORIT	Y APP				•	•	•			998-	9569	1 P	Ρ	19980	0807	
									WO 1	999-1	US17	444	W	19990	0804	

GΙ

Ι

AB An improved process for prepg. the compd. of formula (I) is disclosed which utilizes certain amino acids and divalent cations such as Ni, Co, and Zn to increase titer and decrease the amt. of structural analogs.

IT 120692-19-5P, Pneumocandin A0
RL: BAC (Biological activity or effector, except adverse); BMF
(Bioindustrial manufacture); BPN (Biosynthetic preparation); BSU
(Biological study, unclassified); BIOL (Biological study); PREP
(Preparation)

(prodn. of antibiotic agents by Glarea)

RN 120692-19-5 HCAPLUS

CN Pneumocandin A0 (9CI) (CA INDEX NAME)

Currently available stereo shown.

PAGE 1-A

IT 7440-02-0, Nickel, biological studies
RL: BUU (Biological use, unclassified); BIOL (Biological study);
USES (Uses)

(prodn. of antibiotic pneumocandin derivs. with Glarea lozoyensis)

RN 7440-02-0 HCAPLUS

CN Nickel (8CI, 9CI) (CA INDEX NAME)

Ni

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L41 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1999:708788 HCAPLUS

DOCUMENT NUMBER:

131:322923

TITLE:

A process for the conversion of echinocandin class of peptides to their C4-homotyrosine

monodeoxy analogs

INVENTOR(S):

Mukhopadhyay, Triptikumar; Jayvanti, Kenia;

Kumar, Erra Koteswara Satya Vijaya

PATENT ASSIGNEE(S):

Hoechst Marion Roussel Deutschland GmbH, Germany

SOURCE:

PCT Int. Appl., 18 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	. I	KIND	DATE		APPLI	CATION 1	10.	DATE		
WO 9955727						99-EP27	15	19990	1422	
			AZ, BA,							
DE	, DK, E	E, ES,	FI, GB,	GD,	GE, GH,	GM, HR	HU,	ID,	IL,	IN,
			KP, KR,							
			MX, NO,							
			TR, TT,							
			MD, RU,							
			MW, SD,			ZW, AT	BE,	CH,	CY,	DE,
			GB, GR,							
			GA, GN,							
CA 2327474		AA	19991104		CA 19	99-2327	174	19990	422	
AU 9937096		A1	19991116		AU 19	99-3709	5	19990)422	
BR 9909853		Α :	20001219		BR 19	99-9853		19990	422	
EP 1073675										
R: AT	, BE, C	H, DE,	DK, ES,	FR,	GB, GR,	IT, LI	LU,	NL,	SE,	PT,
IE	, SI, L'	r, LV,	FI, RO							
JP 2002513	033	T2 :	20020508		JP 20	00-5458	35	19990	422	
NO 2000005	258	Α .	20001019		NO 20	000-5258		20001	.019	
PRIORITY APPLN.	INFO.:				EP 1998-	107397	Α	19980	423	
					WO 1999-)422	
OTHER SOURCE(S)	:	CAS	REACT 13	1:32	2923; MF	ARPAT 13	1:322	2923		

Echinocandin type peptides I (X = OH; W, Y, Z = OH, H; R = Me, AΒ

I

Searcher

Shears

308-4994

09/673836

CH2CONH2, H; R' = linoleoyl, 10,12-dimethylmyristoyl, 12-methyltetradecanoyl) were converted to their C4-homotyrosine (C4-htyr) monodeoxy analogs I (X = H) via a single step selective redn. of the C4-htyr hydroxyl group of echinocandins to their monodeoxy analogs under neutral conditions without prior protection/deprotection of the equally facile C5-Orn (ornithine) hydroxyl group and purifn. of the monodeoxy compd. from the crude reaction mixt. Thus, a mixt. of mulundocandin and Raney nickel in a pH 7 ethanol soln. was stirred for 3 h at room temp. to afford 30% deoxymulundocandin, following purifn. by liq.-liq. chromatog. 71018-12-7P, Echinocandin c 138626-63-8P, Deoxymulundocandin 144476-69-7P, Deoxypneumocandin A2 248281-21-2P, Deoxypneumocandin A0 248281-23-4P, Deoxypneumocandin Al RL: SPN (Synthetic preparation); PREP (Preparation) (process for conversion of echinocandin class of peptides to their C4-homotyrosine monodeoxy analogs) 71018-12-7 HCAPLUS

PAGE 1-A

Me-
$$(CH_2)_4$$
 - CH = CH - CH

(CA INDEX NAME)

PAGE 2-A

0

138626-63-8 HCAPLUS

RN

Echinocandin C (9CI)

ΙT

RN CN

CN Deoxymulundocandin (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A | | | Me O

RN 144476-69-7 HCAPLUS
CN Pneumocandin A2, 4-[4-(4-hydroxyphenyl)-L-threonine]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 248281-21-2 HCAPLUS
CN Pneumocandin A0, 4-[4-(4-hydroxyphenyl)-L-threonine]- (9CI) (CA INDEX NAME)

PAGE 1-A

RN 248281-23-4 HCAPLUS
CN Pneumocandin A0, 4-[(.alpha.S)-.alpha.-amino-4-hydroxybenzenebutanoic acid]- (9CI) (CA INDEX NAME)

Ме

PAGE 1-A

REFERENCE COUNT: -- 6 - THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

TIEST REGISTRY' ENTERED AT 12:41:37 ON 17 OCT 2002

L42

7 SEA FILE=REGISTRY ABB=ON PLU=ON (120692-19-5/BI OR 138626-63-8/BI OR 144476-69-7/BI OR 248281-21-2/BI OR 248281-23-4/BI OR 71018-12-7/BI OR 7440-02-0/BI)

FIRE "CAOED' ENTERED AT 12:41:53 ON 17 OCT 2002 L43 0 S L42

FILE 'USPACEULL' ENTERED AT 12:42:01 ON 17 OCT 2002)

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1300 SEA ABB=ON PLU=ON L42/P
O SEA ABB=ON PLU=ON L44 AND (C4(W)(HTYR OR (H OR
L4^{4}
L45
                                                                    HOMO) (W) (TYR OR TYROSINE)))
L50
                                               1288 S L44(S)(L39 OR RANEY(W)(NICKEL OR NI))
                                                            0 S L50(S)(REDUC? OR RED#)
L51
                   (FILE: *CASREAGE ENTERED AT 12:47:10 ON 17 OCT 2002)
L22
                                                                    STR
                                                                                                                                                                                                                          61
                                                                                                                                                                                                                           0
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                                                                         0 22
                                                                                                    G1 19
                                                 OH 23
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                                                                                                                                                                                        OH
                         HO
                 56
                                                                                                                              34 Q
                                     38 NH
                                                                                                                                                                                                     CH3
                                                                                                41
                                                                                                                                                                                                                  36
                                                                                                                                        32
                                                                                                                                        27
                                     43 O
                                                               40
                                                                                                 42
                                        53 G1
                                      54 G1
                                                                                          OH
                                                                        51
                                                                                            52
VAR G1=H/OH
VAR G2=H/CH3/58
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT
                                 IS UNS AT 51
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS
 STEREO ATTRIBUTES: NONE
                                                   14 SEA FIEE CASREACT SSS FUL L22 (
                                                                                                                                                                                                                      217 REACTIONS)
                                                                                                                                                                                                                                                                                        14 DOCS
                                                                 391 VERIFIED
                                                                                                                                         217 HIT RXNS
 100.0% DONE
SEARCH TIME: 00.00.01
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09/673836

L53 ANSWER 1 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

134:296085 CASREACT

TITLE:

FR131535, a novel water-soluble

echinocandin-like lipopeptide: synthesis and

biological properties

AUTHOR(S):

Fujie, A.; Iwamoto, T.; Sato, B.; Muramatsu, H.;

Kasahara, C.; Furuta, T.; Hori, Y.; Hino, M.;

Hashimoto, S.

CORPORATE SOURCE:

Exploratory Research Laboratories, Fujisawa Pharmaceutical Co., Ltd., Ibaraki, Tsukuba-shi,

300-2698, Japan

SOURCE:

Bioorganic & Medicinal Chemistry Letters (2001),

11(3), 399-402

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB The synthesis and biol. properties of a novel water-sol. echinocandin-like lipopeptide, FR131535, are described. This compd. displayed potent in vitro and in vivo antifungal activities. The hemolytic activity of FR901379 was reduced by replacing the acyl side chain. This compd. showed good water-soly., comparable to the natural product FR901379. The synthesis and biol. properties of a novel water-sol. echinocandin-like lipopeptide FR131535 are described.

RX(1) OF 10 **A** ===> B...

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
PAGE 2-A

Na

Α

(1)

В

RX(1) RCT A 138328-74-2

> PRO B 334541-91-2

NTE literature prepn.

REFERENCE COUNT:

THERE ARE 17 CITED REFERENCES AVAILABLE 17

FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

CASREACT COPYRIGHT 2002 ACS L53 ANSWER 2 OF 14

ACCESSION NUMBER:

TITLE:

134:131818 CASREACT

Preparation of novel cyclohexapeptides based on

mulundocandin for use as antifungal agents

INVENTOR(S):

Bansi, Lal; Vitthal, Genbhau Gund; Ashok, Kumar

Gangopadhyay

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland G.m.b.H., Germany

SOURCE:

PCT Int. Appl., 67 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION: - -

PATEN	T NO.		KI	ND	DATE			A.	PPLI	CATI	и ис	o. 1	DATE		
WO 20	010074	168	Α	2	2001	0201		M	0 20	00-E	P676	9 :	2000)715	
WO 20	010074	168	Α	3	2001	1108									
V	V: AE,														
		EE,													
		LT,													
	TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,
	TJ,	TM													

Shears 308-4994 Searcher :

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG EP 1204677 A2 20020515 EP 2000-953050 20000715 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL PRIORITY APPLN. INFO.:

EP 1999-114649 19990727 WO 2000-EP6769 20000715

OTHER SOURCE(S):

MARPAT 134:131818

_R8

OH

Cyclohexapeptides I [R' = alkyl, alkenyl, Ph, biphenyl, terphenyl, naphthyl, alkyl-, alkenyl-, or alkoxyphenyl, linoleoyl, palmitoyl, 12-methylmyristoyl, 10,12-dimethylmyristoyl, or COC6H4OC8H17-p; R1, R3 = OH, CN, CH2NH2, N3, (un) substituted aryl or heterocyclyl with 1-3 of the same or different heteroatoms, aminoalkylamino, (un) substituted alkoxy, etc.; R2, R4 = H, OH; R5 = H, Me; R6 = H, Me, CH2CONH2; R7 = H, Me, OH; R8, R9 = H or secondary aminomethyl] or their pharmaceutically acceptable salts were prepd. for use as antifungal agents. Thus, mulundocandin underwent mono- and dibenzylation on treatment with benzyl alc. and a catalytic amt. of p-toluenesulfonic acid in 1,4-dioxane. Ornithine-5-benzylmulundocandin underwent Mannich reaction with a various secondary amines.

RX(1) OF 69 2 **A** + 3 B ===> C + D...

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

	Me		0			•				
A										
* STRUCTURE	DIAGRAM	TOO LA	RGE FOR	DISPLAY	-	AVAILABLE		FLINE GE 2-A	PRINT	*
	 Me		 							
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HO * Ph	(1)									
3 B										
* STRUCTURE	DIAGRAM	TOO LA	RGE FOR	DISPLAY	-	AVAILABLE		FLINE GE 2-A	PRINT	*
				0		 Me				,
C YIELD 67%						·				
* STRUCTURE	DIAGRAM	TOO LA	RGE FOR	DISPLAY	_	AVAILABLE		FLINE GE 2-A	PRINT	*
D				0		Ме				
YIELD 13%					-			- (
RX (1) RC	CT A 108	351-20	9-8, B 1	00-51-6						
		104-15	-4 TsOH -1 Diox							
		E 144-	-55-8 Na: .8-5 Wat							
		Se	earcher	• 9	Shea	ars 308	3-4994			

09/673836

PRO C 321660-96-2, D 321745-36-2

L53 ANSWER 3 OF 14 CASREACT COPYRIGHT 2002 ACS 134:17732 CASREACT ACCESSION NUMBER: Novel echinocandin derivatives, method for TITLE: preparing same and use as antifungal agents Corbier, Alain; Fauveau, Patrick; INVENTOR(S): Pietre-Dischamp, Nathalie; Schio, Laurent; Vicat, Pascale Hoechst Marion Roussel, Fr. PATENT ASSIGNEE(S):

PCT Int. Appl., 34 pp. SOURCE:

CODEN: PIXXD2 Patent DOCUMENT TYPE:

French LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	TENT 1	NO.		KII	ND	DATE			A	PPLI	CATI	N NC	٥.	DATE		
WO	2000	0751	78	A.	1	2000	1214		M(200	00-F	R156:	9	20000	3608	
	W:	AE,	AG,	AL,	AU,	BA,	BB,	BG,	BR,	CA,	CN,	CR,	CU,	CZ,	DM,	DZ,
														LC,		
														SG,		
														ΚZ,		
		ТJ,					-									
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,
														NL,		
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG
FR	2794	747		A.	1	2000	1215		F	R 199	99-7	252		19990	0609	
EP	1189	932		A.	1	2002	0327		E	P 200	00-9	4045	6	20000	3608	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,
		PT,	IE,	SI,	LT,	LV,	FI,	RO								
PRIORIT	Y APP	LN.	INFO	. :					Fl	R 199	99-72	252		19990	0609	
									W	200	00-F	R156	9	2000	0608	
OTHER SO	OURCE	(S):			MAR	PAT	134:	1773	2							

GΙ

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The invention concerns cyclic peptides I wherein: R = chain contg. AB up to 30 carbon atoms, optionally contg. one or several heteroatoms, one or several heterocycles; either R1 and R2 = H, OH, alkyl optionally substituted, or NR1 forms with the carbon bearing NR1R2 a double bond and R2 is XRa, X being O, NH or N-alkyl and Ra being H, alkyl optionally substituted; R3 = H, OH, CH3; R4 = H, OH; T = H, CH3, CH2CONH2, CH2CN, (CH2)2NH2; Y = H, OH, halogen, OSO3H; W = H, OH; Z = H or CH3. The products of formula I have antifungal properties. Thus, trans-1-[4-[(2-aminocyclo-hexyl)amino]-N2-[[4-[5-[4-(pentyloxy)phenyl]-3-isoxazolyl]phenyl]carbonyl]-L-ornithine]-4-[4-(4-hydroxyphenyl)-L-threonine]-5-L-serine-echinocandin B trifluoroacetate was prepd. and tested for its inhibition of glucan synthase of Candida albicans.

308-4994 Searcher : Shears

RX(1) OF 28 ...2 A + 2 B + 2 C ===> **D** + E

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT * PAGE 2-A

2 A

O HO-C-CH3
$$H_2N$$
 H_2N H_2

D: CM 1

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

D: CM 2

E: CM 1

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

E: CM 2

RX(1) RCT A 310459-17-7, B 38734-69-9

STAGE(1)

RGT F 25895-60-7 NaBH3CN SOL 67-56-1 MeOH

STAGE (2)

RCT C 76-05-1

SOL 7732-18-5 Water, 75-05-8 MeCN

PRO D **310459-08-6**, E 310459-11-1

5

NTE 4A mol. sieves; last step semi-preparative HPLC

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L53 ANSWER 4 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

134:17731 CASREACT

TITLE:

Echinocandin derivatives, method for preparing

same and application as glucan synthase

inhibitors and antifungal agents

INVENTOR(S):

Fauveau, Patrick; Hawser, Stephen; Lebourg,

Gilles; Schio, Laurent

PATENT ASSIGNEE(S):

Hoechst Marion Roussel, Fr.

SOURCE:

PCT Int. Appl., 24 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

GI

French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P	ATENT 1	NO.		KII	ND I	DATE			A	PPLI	CATIO	ои ис	Э.	DATE		
	2000					2000	1014			200		0156	 o	2000	1609	
W	2000															
	W:													CZ,		
		EE,	GD,	GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚP,	KR,	LC,	LK,	LR,
		LT,	LV,	MA,	MG,	MK,	MN,	MX,	ΜZ,	NO,	NΖ,	PL,	RO,	SG,	SI,	SK,
		TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,
		ТJ,	MT													
	RW:													AT,		
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,
														SN,		ΤG
	R 2794													1999		
E	2 1189	933		A:	1 :	2002	0327		E	P 20	00-9	4216	9	2000	0608	
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,
		PT,	ΙE,	SI,	LT,	LV,	FI,	RO								
PRIORI'	TY APP	LN.	INFO	. :					Fl	R 19	99-7	251		1999	0609	
								-	W	20t	00-F	R156	8 .	2000	0608	
OTHER :	SOURCE	(S):			MAR	PAT	134:	1773	1							

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The invention concerns in all possible isomeric forms as well as their mixts., cyclic peptides I wherein: R represents a linear,

branched or cyclic chain; either R1 represents H or CH3 and R2 represents cyclohexyl substituted by an amine, cyanoalkyl; or R1 and R2 form with the nitrogen which bears them a cycle with 3, 4 or 5 carbons optionally substituted by an amine; R3 represents hydrogen, Me or hydroxyl; R4 represents hydrogen or hydroxyl; T represents hydrogen, Me, CH2CONH2, CH2CN, a (CH2)2NH2 or (CH2)2Nalk+X- radical, X being halogen and alk an alkyl radical; Y represents hydrogen, hydroxyl, halogen or OSO3H; W represents H or OH; Z represents H, CH3. The compds. of formula I have antifungal properties. Thus,. Trans 1-[4-[(2-aminocyclohexyl)amino]-N2-[[4''-(pentyloxy)[1,1':4',1''terphenyl]-4-yl]carbonyl]-L-ornithine]-4-[4-(4-hydroxyphenyl)-L-threonine]-5-L-serine-echinocandine B trifluoroacetate was prepd. and tested for its inhibition of glucan synthase of Candida albicans and of the enzyme prepd. from Aspergillus fumigatus.

$$RX(1)$$
 OF 12 ...2 A + 2 B + 2 C ===> **D** + E

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

2 A

Me F F C CO₂H F C CO₂H

2 B 2 C
$$(1)$$

D: CM 1

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

D: CM 2

E: CM 1

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

E: CM 2

RX(1) RCT A 227472-55-1, B 19777-66-3

STAGE(1)

RGT F 121-44-8 Et3N SOL 67-56-1 MeOH

STAGE (2)

RGT G 25895-60-7 NaBH3CN

STAGE (3)

RCT C 76-05-1

SOL 7732-18-5 Water, 75-05-8 MeCN

RO D **310461-86-0**, E 310461-89-3

NTE 1st stage siliporite grains; last stage semi-preparative

HPLC

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L53 ANSWER 5 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

132:308664 CASREACT

TITLE:

Photochemical process for conversion of the 1,2-diol moiety of an echinocandin compound to

the 1-deoxy-2-keto analog

INVENTOR(S):

Hitchcock, Stephen Andrew; Gregory, George

Stuart

PATENT ASSIGNEE(S):

Eli Lilly and Company, USA

SOURCE:

PCT Int. Appl., 28 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.	KI	ND I	DATE			A	PPLI	CATI	N NC	o. !	DATE		
		- -												
WO 2000	024694	4 A:	1 :	2000	0504		W	0 19	99 - U	S253	01	1999:	1027	
W:	AE, A	AL, AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,
		CZ, DE,												
		IL, IN,												
		JV, MA,												
	SD, S	SE, SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,
		YU, ZA,												
RW:		SM, KE,											CH,	CY,
		OK, ES,												
	BJ, C	CF, CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	ΤG	
PRIORITY APP	LN. IN	NFO.:					U	s 19	98-1	0593	6P	1998:	1028	
OTHER SOURCE	(S):		MAR	PAT	132:	3086	64							

AB A method for converting an epoxy or hydroxy moiety to a 1-deoxy-2-keto moiety is described which includes: (1) reacting a compd. having an epoxy or hydroxy moiety with a thiophenol and (2) irradiating the 1-phenylthio-2-hydroxy moiety with UV or near-UV radiation to convert the 1-phenylsulfide-2-hydroxy moiety to a 1-deoxy-2-keto moiety. The process was used to modify the cyclic peptide ring system of an echinocandin-type compd. contg. a 1,2-diol moiety to produce new keto analogs.

RX(1) OF 3 **A** + B ===> C...

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

$$\begin{array}{c} I \\ S \\ \star \\ H \end{array}$$

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Searcher : Shears

308-4994

С

RCT A 166663-25-8, B 37972-89-7 RX(1)

PRO C 266317-25-3

SOL 75-05-8 MeCN, 67-56-1 MeOH 4

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 6 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

TITLE:

131:322923 CASREACT

A process for the conversion of echinocandin class of peptides to their C4-homotyrosine

monodeoxy analogs

INVENTOR(S):

Mukhopadhyay, Triptikumar; Jayvanti, Kenia;

Kumar, Erra Koteswara Satya Vijaya

PATENT ASSIGNEE(S): SOURCE:

Hoechst Marion Roussel Deutschland GmbH, Germany

PCT Int. Appl., 18 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

						- • •			-							
PA.	rent :	NO.		KI	ND I	DATE			A	PPLI	CATI	ои ис	o.	DATE		
WO	9955	 727		A	1	1999	1'104		W	0 19	99-E	P271	5	19990	0422	
	W:	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,
														ID,		
		IS,	JP,	ΚĖ,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,
		MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,
		SK,	SL,	TJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,
		AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM							
	RW:	GH,	GM,	ΚE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,

308-4994 Searcher : Shears

09/673836

DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG 19990422 19991104 CA 1999-2327474 CA 2327474 AAAU 1999-37096 19990422 AU 9937096 Α1 19991116 BR 9909853 20001219 BR 1999-9853 19990422 Α 20010207 EP 1999-919261 19990422 EP 1073675 A1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, SI, LT, LV, FI, RO JP 2000-545885 19990422 JP 2002513033 T2 20020508 NO 2000005258 Α 20001019 NO 2000-5258 20001019 EP 1998-107397 19980423 PRIORITY APPLN. INFO.: WO 1999-EP2715 19990422

OTHER SOURCE(S):

MARPAT 131:322923

GΙ

... Š

Echinocandin type peptides I (X = OH; W, Y, Z = OH, H; R = Me, AΒ CH2CONH2, H; R' = linoleoyl, 10,12-dimethylmyristoyl, 12-methyltetradecanoyl) were converted to their C4-homotyrosine (C4-htyr) monodeoxy analogs I (X = H) via a single step selective redn. of the C4-htyr hydroxyl group of echinocandins to their monodeoxy analogs under neutral conditions without prior protection/deprotection of the equally facile C5-Orn (ornithine) hydroxyl group and purifn. of the monodeoxy compd. from the crude reaction mixt. Thus, a mixt. of mulundocandin and Raney nickel in a pH 7 ethanol soln. was stirred for 3 h at room temp. to afford 30% deoxymulundocandin, following purifn. by liq.-liq. chromatog.

Ι

RX(1) OF 1

STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT * PAGE 2-A

> 1 0

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308-4994 Searcher Shears

(1)

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

0

B YIELD 75%

RX(1) RCT A 54651-05-7

RGT C 7440-02-0 Ni PRO B 71018-12-7

SOL 64-17-5 EtOH

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L53 ANSWER 7 OF 14 CASREACT COPYRIGHT 2002 ACS

6

ACCESSION NUMBER:

129:276286 CASREACT

TITLE: AUTHOR(S):

Studies on the phosphorylation of LY303366 Udodong, Uko E.; Turner, William W.; Astelford, Bret A.; Brown, Frank, Jr.; Clayton, Marcella

T.; Dunlap, Steven E.; Frank, Scott A.; Grutsch, John L.; LaGrandeur, Lisa M.; Verral, Daniel E.;

Werner, John A.

CORPORATE SOURCE:

Lilly Research Laboratories, Lilly Corporate

Center, Indianapolis, IN, 46285, USA

SOURCE:

Tetrahedron Letters (1998), 39(34), 6115-6118

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Phosphorylation of LY303366 was studied in THF and DMF. Benzyl phosphate (I) could be prepd. in excellent yield using LiOH as the base. Both I and the derived phosphonic acid monosodium salt were prone to undergo hydrolytic dephosphorylation.

RX(1) OF 1 **A** + B ===> C

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

Α

$$\begin{array}{c|cccc}
Ph & O & O & Ph \\
Ph & O & P & O \\
Ph & O & Ph \\
B & & & & & \\
\end{array}$$

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

YIELD 33%

RCT A 166663-25-8, B 990-91-0 RX(1)

RGT D 1310-65-2 LiOH PRO C 213669-65-9

SOL 75-09-2 CH2Cl2, 109-99-9 THF

6

THERE ARE 6 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L53 ANSWER 8 OF 14 CASREACT COPYRIGHT 2002 ACS

126:212437 CASREACT ACCESSION NUMBER:

Preparation of cyclic peptide antifungal agents TITLE:

Rodriguez, Michael John INVENTOR(S): Lilly, Eli, and Co., USA PATENT ASSIGNEE(S):

Eur. Pat. Appl., 22 pp. SOURCE: CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

		-															
	PAT	ENT	NO.	:	KIN	ND ,	DATE			Al	PPLIC	CATI	ои ис	Э.	DATÉ		
	EΡ	7570	58 -	•	. A1		1997	0205	. 8	E1	P 199	96-30	05345	5 ·	1996	0722	
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		~	PT,	SE				,									
	110		•	~ —	70		1997	ΛE 1 2		T10	s 199	95-51	06790	Ω	19950	1725	
	US	5629	289		Α		1997	0212						-			
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	とり	2151	030		1.)	2001	OTOI									
	WO	9705	163		A1	l	1997	0213		W	0 199	96-U	S121:	11	1996	3723	
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•			ıs,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LK,	LR,	LS,	LT,	LV,	MD,	MG,	MK,
			MN.	MW.	MX.	NO.	NZ,	PL.	RO,	RU.	SD.	SG,	SI,	SK,	ТJ,	TM,	TR,

09/673836

TT, UA, UG, US, UZ
RW: KE, LS, MW, SD, SZ, UG, BF, BJ, CF, CG, CI, CM, GA, GN, ML,
MR, NE, SN, TD, TG

AU 9665938 A1 19970226 AU 1996-65938 19960723 JP 11510165 T2 19990907 JP 1996-507687 19960723 PRIORITY APPLN. INFO.: US 1995-506790 19950725

WO 1996-US12111 19960723

OTHER SOURCE(S):

MARPAT 126:212437

GT

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- Provided are pharmaceutical formulations, and methods of inhibiting AΒ fungal and parasitic activity using cyclopeptides I [R11 = H, CH2OH, CHMeOH, CH(OH)CH2CONH2; R12 = H, CH2OH, CHMeOH; R13 = H, Me; R31 = H, OH, OR30; R30 = C1-6 alkyl, PhCH2, (CH2)2SiMe3, CH2CH:CH2, CH2CH(OH)CH2OH, (CH2)aCO2H, (CH2)bNR41R42, (CH2)cPOR43R44, (CH2CH2O)d(C1-6)alkyl; a, b, c = 1-6; R41, R42 = H, C1-6 alkyl; R41R42 = (CH2)e; R43, R44 = OH, C1-6 alkoxy; d = 1, 2; e = 3-5; R32,R21, R22, R23, R24 = OH, H; R0 = OH, OPO3H2, OP(O)(OH)R1, OP(O)(OH)OR1, R1 = C1-6 alkyl, Ph, p-halophenyl, p-nitrophenyl, PhCH2, p-halobenzyl, p-nitrobenzyl; R2 = COC6H4R3; R3 = C6H4R5-4, C.tplbond.CC6H4R6-4, p-C6H4C.tplbond.CC6H4R7-4, p-C6H4C6H4R8-4; R5, R6, R7, R8 = H, C1-12 alkyl, C2-12 alkynyl, C1-12 alkoxy, C1-12alkylthio, halo, O(CH2)m[O(CH2)n]pO(C1-12 alkyl), O(CH2)qXR4; m =2-4; n = 2-4; p = 0, 1; q = 2-4; X = pyrrolidino, piperidino, piperazino; R4 = H, C1-12 alkyl, C3-12 cycloalkyl, benzyl, C3-12 cycloalkylmethyl; with the proviso that at least 1 of R11 and R12 must be H] or pharmaceutically acceptable salt thereof. Thus, acylation of 348.1 g (60.2 mmol) antibiotic A 30912A nucleus with 26.0 g (48.2 mmol) terphenyl active ester Me(CH2)40-p-C6H4-p-C6H4-p-C6H4CO2C6H2Cl3-2,4,5 in 8.5 L of DMF gave 18 g acylated deriv. II (R11 = R12 = CHMeOH, R31 = R32 = OH) (III). Treatment of 5 g III with 17 mL CF3CO2H and 35 mL Et3SiH in 250 mL CH2Cl2 gave 3.872 g (79%) reduced deriv. II (R11 = R12 = CHMeOH, R31 = R32 = H), which underwent retro-aldol condensation by treatment with 2.51 g (22.6 mmol) Me3N+O- in 20 mL of a 1:1 mixt. of MeCN and DMF at 100.degree. for 24 h to give 72% II (R11 = R12 = R31 = R32 = H). Pharmaceutical formulations contg. II (R11 = R12 = R31 = R32 = H) arte given.

RX(1) OF 6 A + B ===> C..

Α

В

(1)

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Searcher :

Shears

308-4994

С

RCT A 79411-15-7, B 158937-65-6 RX(1) PRO C 166663-25-8 SOL 68-12-2 DMF

L53 ANSWER 9 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

125:115162 CASREACT

TITLE:

Process for performing retro-aldol reactions

using amine oxide agents Rodriguez, Michael John

INVENTOR(S):

Lilly, Eli, and Co., USA PCT Int. Appl., 19 pp.

PATENT ASSIGNEE(S): SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	rent :	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	٥.	DATE		
WO	9615	 142		 A	 1	1996	0523		W	0 19	95-U	S146	13	1995	1113	
														DE,		EE,
														LK,		
														PT,		
		SD,	SE,	SG,	SI,	SK										
•	RW:	KE,														
		ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,
						TD,										
	2205															
	9641															
EΡ	7871															
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LI,	LU,	NL,	PT,
		SE														
J.P-	-1050	8852				1998					95-5			1995		
US	5677	423		Α		1997	1014		Ü	S 19	96-7	6358	4	1996	1210	
	_	/														

308-4994 Searcher : Shears

09/673836

Ι

PRIORITY APPLN. INFO.:

US 1994-339525 19941115 WO 1995-US14613 19951113

OTHER SOURCE(S):

MARPAT 125:115162

GI

A process for removing .beta.-hydroxy groups from AΒ .beta.-hydroxy-contg. compds. id disclosed. The process involves the use of a retro-aldol-promoting reagent selected from the group consisting of trimethylamine-N-oxide, triethylamine-N-oxide, trimethylamine-N-oxide hydrate, and trimethylamine-hydrate and requires dissoln. of the substrate in an aprotic solvent and reaction under elevated temps. The process is broadly applicable to a variety of substrates including complex cyclic peptides, linear peptides, and nonpeptides. Thus, 0.25~g cyclopeptide R106-1 (I; R = CMe2OH), obtained by fermn. from Aureobasidium pullulans, was dissolved in 2.5 mL MeCN and 0.25 g trimethylamine N-oxide hydrate added all at once. The reaction mixt. was heated at 70.degree. for 24 h, cooled to room temp., concd. under vacuum, dissolved in EtOAc, washed with cold 10% HCl, satd. NaHCO3, and brine, and purified by reverse-phase preparative HPLC to yield 0.22 g (92%) of sarcosine-contg. cyclopeptide I (R = H).

RX(2) OF 3 $\mathbf{E} ===> \mathbf{F}$

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT_*

Ε

(2)

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

F

RX(2) RCT E 179118-65-1

RGT C 136449-78-0 Methanamine, N, N-dimethyl-, N-oxide,

monohydrate

PRO F 179118-66-2

SOL 75-05-8 MeCN, 68-12-2 DMF

NTE regioselective

CASREACT COPYRIGHT 2002 ACS L53 ANSWER 10 OF 14

ACCESSION NUMBER:

117:234513 CASREACT

TITLE:

Reduction studies of antifungal echinocandin

breezery

lipopeptides. One step conversion of

echinocandin B to echinocandin C

AUTHOR(S): CORPORATE SOURCE: Balkovec, James M.; Black, Regina M. Dep. Synth. Chem. Res., Merck Res. Lab.,

Rathway, NJ, 07065-0900, USA

SOURCE:

Tetrahedron Letters (1992), 33(32), 4259-32

CODEN: TELEAY; ISSN: 0040-4039

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Sodium cyanoborohydride in trifluoroacetic acid selectively reduced AΒ the C5-hydroxyornithine and C4-hydroxyhomotyrosine carbinols to methylene groups in echinocandin lipopeptides. The selective redn. of either hydroxyl is also described. The first conversion of echinocandin B to echinocandin C was accomplished.

RX(3) OF 12 **F** + G ===> A...

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT * PAGE 2-A

> 0

F

$$O_2N$$
 O_2N
 O_2N

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

308-4994 Shears Searcher :

Α

RCT F **54651-06-8**, G 5070-13-3 RX (3)

RGT H 1310-65-2 LiOH PRO A 144448-04-4 SOL 872-50-4 NMEP

CASREACT COPYRIGHT 2002 ACS L53 ANSWER 11 OF 14

ACCESSION NUMBER:

117:70296 CASREACT

TITLE:

Preparation and structure-activity relationships

of simplified analogs of the antifungal agent

cilofungin: a total synthesis approach

AUTHOR (S):

Zambias, Robert A.; Hammond, Milton L.; Heck, James V.; Bartizal, Ken; Trainor, Charlotte; Abruzzo, George; Schmatz, Dennis M.; Nollstadt,

Karl M.

CORPORATE SOURCE:

SOURCE:

Merck Res. Lab., Rahway, NJ, 07065, USA

Journal of Medicinal Chemistry (1992), 35(15),

2843-55

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

LANGUAGE:

Journal English

GI

308-4994 Searcher Shears

The echinocandins are a well-known class of lipopeptides AB characterized by their potent antifungal activity against Candida species. The mechanism of action of the echinocandins is generally thought to be the inhibition of .beta.-1,3-glucan synthesis, an important structural component in the cell wall of Candida species. Extensive structure-activity studies on the fatty acid side chain of echinocandin B led to the prepn. of the clin. candidate cilofungin. We now report the prepn., by solid-phase synthesis, of a series of simplified analogs of cilofungin in which the unusual amino acids found in the echinocandins were replaced with more readily accessible natural amino acids. The solid-phase approach to the total synthesis of these analogs allowed us to conveniently explore structural modifications that could not be accomplished by chem. modification of the natural product. The simplest analog I [R =p-[Me(CH2)70]C6H4CONH] showed no biol. activity. Structural complexity was then returned to the system in a systematic fashion so as to reapproach the original cilofungin structure. Antifungal activity and the inhibition of .beta.-1,3-glucan synthesis were monitored at each step of the process, thereby revealing the basic structure-activity relationships of the amino acids and the minimal structural requirements for biol. activity in the echinocandin ring system. The results suggests that the 3-hydroxy-4-methylproline residue enhances activity but the L-homotyrosine residue is crucial for both antifungal activity and the inhibition of .beta.-1,3-glucan synthesis.

Ι

RX(11) OF 32 ...AE ===> AH

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

PAGE 2-A

ΑE

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT * PAGE 2-A

(CH₂)₇-Me

AΗ YIELD 16%

RCT AE 141806-24-8 RX(11)

C 26386-88-9 (PhO)2P(O)N3, D 144-55-8 NaHCO3 RGT

PRO AH 141806-25-9 SOL 68-12-2 DMF NTE Key step

L53 ANSWER 12 OF 14 CASREACT COPYRIGHT 2002 ACS

108:38346 CASREACT ACCESSION NUMBER:

TITLE: Mulundocandin, a new lipopeptide antibiotic.

II. Structure elucidation

Mukhopadhyay, Triptikumar; Ganguli, B. N.; Fehlhaber, H. W.; Kogler, H.; Vertesy, L. Res. Cent., Hoechst India Ltd., Bombay, 400 080, AUTHOR(S):

CORPORATE SOURCE:

India

J. Antibiot, (1987), 40(3), 281-9 SOURCE:

CODEN: JANTAJ; ISSN: 0021-8820

DOCUMENT TYPE: Journal

LANGUAGE: English

GΙ

Ι

AB Mulundocandin, a new antifungal antibiotic, was shown to have structure I by high field NMR expts., e.g., homo- and heteronuclear correlation spectra, distortionless enhancement by polarization transfer (DEPT) spectra as well as nuclear Overhauser effect. The compd. is a lipopeptide belonging to the echinocandin class.

RX(1) OF 5 **A** ===> B...

Α

RX(1) RCT A 108351-20-8 RGT C 7647-01-0 HC1 PRO B 5502-94-3 SOL 7732-18-5 Water

L53 ANSWER 13 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

107:237279 CASREACT

TITLE:

Synthesis of the cyclic hexapeptide echinocandin

D. New approaches to the asymmetric synthesis

of .beta.-hydroxy .alpha.-amino acids

AUTHOR(S):

Evans, David A.; Weber, Ann E.

CORPORATE SOURCE:

Dep. Chem., Harvard Univ., Cambridge, MA, 02138,

USĀ

SOURCE:

J. Am. Chem. Soc. (1987), 109(23), 7151-7

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The total synthesis of echinocandin D (I, Lin = linoleyl) was achieved using asym. glycine enolate aldol methodol. for the prepn. of 2 constituent .beta.-hydroxy amino acids. Protected hydroxy amino acids II and III were prepd. in 4 steps each from oxazolidinones IV (R = CH2Ph, R1 = H, R2 = NCS) and IV (R = H, R1 = CH2Ph, R2 = Br), resp. In both prepns., asym. aldol addn. was used to establish the abs. stereochem. relationships at both OH and N-bearing asym. centers. II and III were integrated into the synthesis of I.

RX(1) OF 177 ...A ===> B...

$$=$$
 CH $-$ CH $_2-$ CH $=$ CH $-$ (CH $_2$) $_4-$ Me

Α

 $\xrightarrow{(1)}$

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

RX(1) RCT A 104197-62-8

RGT C 26386-88-9 (PhO) 2P(O) N3, D 121-44-8 Et3N

PRO B **71018-13-8** SOL 68-12-2 DMF

L53 ANSWER 14 OF 14 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

105:173030 CASREACT

TITLE:

Total synthesis of echinocandins. II. Total

synthesis of echinocandin D via efficient

peptide coupling reactions

AUTHOR(S):

Kurokawa, Natsuko; Ohfune, Yasufumi

CORPORATE SOURCE:

SOURCE:

Suntory Inst. Bioorg. Res., Osaka, 618, Japan J. Am. Chem. Soc. (1986), 108(19), 6043-5

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

LANGUAGE:

Journal English

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Echinocandin D (I, R = H) was prepd. by deblocking hexapeptide II [R = OMe, R1 = CH2NHCO2CMe3, R2 = H, R3 = Si(CMe3)Me2] (III) and cyclizing the resulting II (R = OH, R1 = CH2NH2, R2 = R3 = H) by diphenylphosphoryl azide. The deblocking of II [R = NH2, R1 = CH(OMe)2, R2 = OSi(CMe3)Me2, R3 = Si(CMe3)Me2] (IV) followed by an attempted cyclization failed to give echinocandin C (I, R = OH). III and IV were prepd. from their amino acid constituents via peptide coupling reactions.

RX(1) OF 188

...A ===> B...

PAGE 1-A

PAGE 1-B

= CH- CH $_2-$ CH= CH- (CH $_2$) 4- Me

Α

 $\xrightarrow{(1)}$

PAGE 1-A

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

RX(1) RCT A 104197-62-8

RGT C 26386-88-9 (PhO) 2P(O) N3

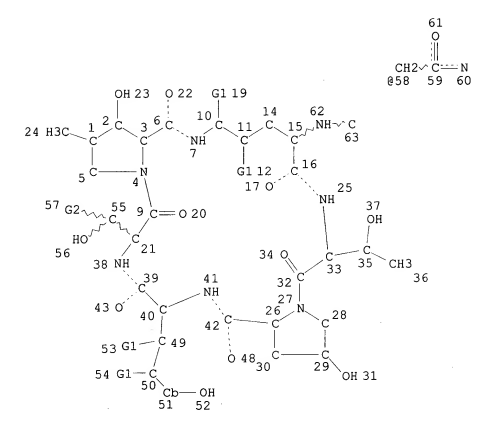
PRO B **71018-13-8**

=> fil djsmds, cheminfo, chemreaet FILE 'DJSMDS' ENTERED AT 12:48:29 ON 17 OCT 2002 COPYRIGHT (C) 2002 THOMSON DERWENT

FILE 'CHEMINFORMRX' ENTERED AT 12:48:29 ON 17 OCT 2002 COPYRIGHT (CAFIZ-CHEMIE BERLIN

FILE 'CHEMREACT' ENTERED AT 12:48:29 ON 17 OCT 2002 COPYREGHT (C) Springer-Verlag/InfoChem GmbH (IC)

=> d que stat; d bib ab fhit L22 STR



VAR G1=H/OH VAR G2=H/CH3/58 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS UNS AT 51 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS

STEREO ATTRIBUTES: NONE

-1 SEA 122

ANSWER 1 OF 1 CHEMREACT COPYRIGHT 2002 SPRINGER/IC L54

AN 1417229 CHEMREACT

DN - 88011612 --

ΑU EVANS DAVID A.; WEBER ANN E.

J. Am. Chem. Soc., 109, 7151-7157 (1987) CODEN: JACSAT ISSN: 0002-7863 SO

LA English

RX(1) OF 1

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Shears 308-4994 Searcher :

TITLE:

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
         RCT A, 164651701
               RCT.STE: S,S,S,S,R,S,R,S,R,S,R,S
          PRO
               B, 71327204
               PRO.STE: S,S,S,S,R,S,R,S,S,S,R,R
          ΥD
               50.0 %
          KW
               IR
                                                                 - Key ferms
claim 2
    FILE REGISTRY ENTERED AT 12:49:24 ON 17 OCT 2002
                E MULUNDOCANDIN/CN 5
              1 SEA ABB=ON PLU=ON MULUNDOCANDIN/CN
L55
                E DEOXYMULUNDOCANDIN/CN 5
              1 SEA ABB=ON PLU=ON DEOXYMULUNDOCANDIN/CN
L56
    FILE LICAPLUS! ENTERED AT 12:49:54 ON 17 OCT 2002
             13 SEA ABB=ON PLU=ON L55 OR MULUNDOCANDIN
L57
              4 SEA ABB=ON PLU=ON L57 AND (L56 OR DEOXYMULUNDOCANDIN
L58
                OR DEOXY MULUNDOCANDIN)
              3 SEA ABB=ON PLU=ON L58 NOT L41
L59
L59 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2002 ACS
                         2002:255768 HCAPLUS
ACCESSION NUMBER:
                         137:201573
DOCUMENT NUMBER:
                         Synthesis of new echinocandin derivatives via a
TITLE:
                         diol-keto transposition
                         Aszodi, Jozsef; Fauveau, Patrick; Melon-Manguer,
AUTHOR(S):
                         Dominique; Ehlers, Eberhard; Schio, Laurent
                         Medicinal Chemistry, Aventis Pharma,
CORPORATE SOURCE:
                         Romainville, F-93235, Fr.
                         Tetrahedron Letters (2002), 43(16), 2953-2956
SOURCE:
                         CODEN: TELEAY; ISSN: 0040-4039
                         Elsevier Science Ltd.
PUBLISHER:
DOCUMENT TYPE:
                         Journal
                         English
LANGUAGE:
     A new diol-carbonyl transposition reaction has been discovered in
AB
     echinocandin type structures. An .alpha.-hydroxy hemiaminal moiety
     has been shown to undergo a pinacol-type rearrangement in the
     presence of trimethylsilyl iodide to afford ketone derivs. Applied
     to deoxymulundocandin, this transposition led to a useful
     intermediate for further chem. modification.
     108351-20-8, Mulundocandin 138626-63-8,
TT
     Deoxymulundocandin
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of in the prepn. of deoxymulundocandin
        derivs. via diol-carbonyl transposition reaction)
                               THERE ARE 23 CITED REFERENCES AVAILABLE
                         23
REFERENCE COUNT:
                               FOR THIS RECORD. ALL CITATIONS AVAILABLE
                               IN THE RE FORMAT
L59 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                         2001:618023 HCAPLUS
DOCUMENT NUMBER:
                         135:180953
                         Preparation of novel echinocandin derivatives as
```

Searcher :

308-4994

Shears

fungicides

INVENTOR(S):

Courtin, Olivier; Dussarat, Arlette;

Melon-Manguer, Dominique; Schio, Laurent

PATENT ASSIGNEE(S): SOURCE:

Aventis Pharma S.A., Fr. PCT Int. Appl., 30 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.				KIND DATE				APPLICATION NO. DATE									
	WO 20010608				 A	20010823				WO 2001-FR419					20010214			
	1	W:	ΑE,	AG,	AL,	AU,	BA,	BB,	BG,	BR,	ΒZ,	CA,	CN,	CR,	CU,	CZ,	DM,	
			DZ,	EE,	GD,	GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚP,	KR,	LC,	LK,	
			LR,	LT,	LV,	MA,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,	SG,	SI,	SK,	
			TT,	UA,	US,	UZ,	VN,	YU,	ZA,	AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	
			TM															
		RW:	GH,	GM,	KE,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	ΤZ,	UG,	ZW,	AT,	BE,	CH,	
			CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,	NL,	PT,	SE,	
			TR,	BF,	BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	
			TG															
FR 2804957 A1 20010817 FR 2000-1844												344	20000215					
PRIORITY APPLN. INFO.: FR 2000-1844													A	2000	3215			
OTHER	R SOU	RCE	(S):			MAR:	PAT :	135:	1809	53								
GT																		

AΒ Echinocandin derivs. I [R1 = H, OH, (un) substituted alkoxy, alkenyloxy or alkynyloxy; R3 = H, Me, OH; R4, W = H, OH; A = O, CH2, NH; B is a steroid residue; T = H, Me, CH2CONH2, CH2C.tplbond.N, (CH2)2NH2 or alkylaminoethyl; Y = H, OH, halo, OSO3H or salts; Z = H, Me] were prepd. as antifungal agents. Thus, 1-[(4R,5R)-4,5dihydroxy-N2-[[[(3.beta., 22E)-ergosta-5,7,22-trien-3-

> 308-4994 Searcher : Shears

yl]oxy]carbonyl]-L-ornithine]deoxymulundocandin was prepd. by treating ergosterol with diphosgene in CH2Cl2 in the presence of Et3N and treating the product with deoxymulundocandin.

IT 108351-20-8, Mulundocandin 138626-63-8,

Deoxymulundocandin

RL: RCT (Reactant); RACT (Reactant or reagent)

3

(prepn. of novel echinocandin derivs. as fungicides)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L59 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1992:527867 HCAPLUS

DOCUMENT NUMBER:

117:127867

TITLE:

Deoxymulundocandin-a new echinocandin

type antifungal antibiotic

AUTHOR(S):

Mukhopadhyay, Triptikumar; Roy, Kirity; Bhat, R. G.; Sawant, S. N.; Blumbach, J.; Ganguli, B. N.;

Fehlhaver, H. W.; Kogler, H.

CORPORATE SOURCE:

Res. Cent., Hoechst India Ltd., Bombay, 400 080,

India

SOURCE:

Journal of Antibiotics (1992), 45(5), 618-23

CODEN: JANTAJ; ISSN: 0021-8820

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GΙ

An ew echinocandin type antifungal antibiotic, deoxymulundocandin (I), C48H77N7O15, was isolated from the culture filtrate and mycelia of a fungal culture, Aspergillus sydowii (Bainier and Sartory) Thom and Church var. nov. mulundensis Roy (Culture No. Y-30462). Its structure was established by comparative GC-MS analyses of the derivatized acid hydrolyzates of deoxymulundocandin and mulundocandin as well as by the high field NMR expts. (COSY, NOESY and DEPT).

IT 138626-63-8, Deoxymulundocandin

RL: BIOL (Biological study)

(antifungal antibiotic, from Aspergillus sydowii)

Searcher: Shears 308-4994

Т

(ELLE MEDITINE, BIOSIS, EMBASE, WPIDS, JICST-EPLUS, JAPIO, CONFSCI, SCISEARCH, GBNB, CIN, CEN, CASREACT, CHEMINFORMRX, CHEMREACT, DJSMDS' ENTERED AT 12:52:08 ON 17 OCT 2002)

7 S L58

3 DUPEREMEL 60 (4 DUPLICATES REMOVED)

L61 ANSWER 1 OF 3 SCISEARCH COPYRIGHT 2002 ISI (R)

ACCESSION NUMBER: 2002:389532 SCISEARCH

THE GENUINE ARTICLE: 545TP

Synthesis of new echinocandin derivatives via a TITLE:

diol-keto transposition

Aszodi J; Fauveau P; Melon-Manguer D; Ehlers E; AUTHOR:

Schio L (Reprint)

Aventis Pharma, Med Chem, 102 Route Noisy, F-93235 CORPORATE SOURCE:

Romainville, France (Reprint); Aventis Pharma, Med Chem, F-93235 Romainville, France; Aventis Pharma, Process Dev Biochem, Biol Sud, D-65956 Frankfurt,

Germany

COUNTRY OF AUTHOR:

France; Germany

SOURCE:

TETRAHEDRON LETTERS, (15 APR 2002) Vol. 43, No. 16,

pp. 2953-2956.

Publisher: PERGAMON-ELSEVIER SCIENCE LTD, THE BOULEVARD, LANGFORD LANE, KIDLINGTON, OXFORD OX5

1GB, ENGLAND. ISSN: 0040-4039. Article; Journal

DOCUMENT TYPE:

LANGUAGE:

English

REFERENCE COUNT:

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS A new diol-carbonyl transposition reaction has been discovered in AΒ echinocandin tape structures. All alpha-hydroxy hemiaminal moiety has been shown to undergo a pinacol-type rearrangement in the presence of trimethylsilyl iodide to afford ketone derivatives, Applied to deoxymulundocandin. this transposition led to a useful intermediate for further chemical modification. (C) 2002 Elsevier Science Ltd. All rights reserved.

L61 ANSWER 2 OF 3 CASREACT COPYRIGHT 2002 ACS

ACCESSION NUMBER:

131:322923 CASREACT

TITLE:

A process for the conversion of echinocandin class of peptides to their C4-homotyrosine

monodeoxy analogs

INVENTOR(S):

Mukhopadhyay, Triptikumar; Jayvanti, Kenia;

Kumar, Erra Koteswara Satya Vijaya

PATENT ASSIGNEE(S):

Hoechst Marion Roussel Deutschland GmbH, Germany

PCT Int. Appl., 18 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE _____ WO 1999-EP2715 19991104 19990422 WO 9955727 A1

W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN,

308-4994 Searcher : Shears

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IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD,
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                     CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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                                            CA 1999-2327474
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                                                              19990422
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                             19991116
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                                                              19990422
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                             20001219
                                            EP 1999-919261
                                                              19990422
     EP 1073675
                       A1
                             20010207
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             IE, SI, LT, LV, FI, RO
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                             20020508
                                            JP 2000-545885
     JP 2002513033
                       T2
                                                              20001019
                                            NO 2000-5258
     NO 2000005258
                             20001019
                                            EP 1998-107397
                                                              19980423
PRIORITY APPLN. INFO.:
                                                              19990422
                                            WO 1999-EP2715
```

OTHER SOURCE(S):

MARPAT 131:322923

Echinocandin type peptides I (X = OH; W, Y, Z = OH, H; R = Me, AΒ CH2CONH2, H; R' = linoleoyl, 10,12-dimethylmyristoyl, 12-methyltetradecanoyl) were converted to their C4-homotyrosine (C4-htyr) monodeoxy analogs I (X = H) via a single step selective redn. of the C4-htyr hydroxyl group of echinocandins to their monodeoxy analogs under neutral conditions without prior protection/deprotection of the equally facile C5-Orn (ornithine) hydroxyl group and purifn. of the monodeoxy compd. from the crude reaction mixt. Thus, a mixt. of mulundocandin and Raney nickel in a pH 7 ethanol soln. was stirred for 3 h at room temp. to afford 30% deoxymulundocandin, following purifn. by liq.-liq. chromatog.

Ι

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

MEDLINE L61 ANSWER 3 OF 3

DUPLICATE 1

ACCESSION NUMBER: 92324937 MEDLINE

> 308-4994 Searcher Shears

DOCUMENT NUMBER:

92324937 PubMed ID: 1624363

TITLE:

Deoxymulundocandin--a new echinocandin type

antifungal antibiotic.

AUTHOR:

Mukhopadhyay T; Roy K; Bhat R G; Sawant S N; Blumbach

J; Ganguli B N; Fehlhaber H W; Kogler H

CORPORATE SOURCE:

Microbiology Department, Hoechst India Limited,

Mulund, Bombay.

SOURCE:

JOURNAL OF ANTIBIOTICS, (1992 May) 45 (5) 618-23.

Journal code: 0151115. ISSN: 0021-8820.

PUB. COUNTRY:

Japan

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199208

ENTRY DATE:

Entered STN: 19920821

Last Updated on STN: 19920821 Entered Medline: 19920813

AB A new echinocandin type antifungal antibiotic,

deoxymulundocandin, C48H77N7O15, was isolated from the culture filtrate and mycelia of a fungal culture, Aspergillus sydowii (Bainier and Sartory) Thom and Church var. nov. mulundensis Roy (Culture No. Y-30462). The structure was established by comparative GC-MS analyses of the derivatized acid hydrolysates of deoxymulundocandin and mulundocandin as well as by the high field NMR experiments (COSY, NOESY and DEPT).

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